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PATENT SPECIFICATION



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COMPLETE SPECIFICATION

Process for the Manufacture of Esters, especially of Ethyl Acetate

We, Usines de Melle, a body corporate organised under the Laws of the Republic of France, of Melle, Deux-Sevres, France (on our own behalf and as Assignees of 5 HENRI MARTIN GUINOT), do hereby declare the nature of this invention and in what manner the same is to be performed, to be particularly described and ascertained in and by the following statement:

This invention relates to the manufacture of the statement of the statement

ture of esters, particularly of ethyl acetate.

It is already known to pass alcohol over dehydrogenation catalysts heated at temdehydrogenation catalysts heated at temperatures between 250° and 500° C. and
kept under a pressure greater than 10
atmospheres with a view to obtaining its
partial conversion into various products
such as ethyl acetate, butyl alcohol,
acetaldehyde acetane crotonyl alcohol. 20 acetaldehyde, acetone, crotonyl alcohol, butyl and crotonic acetates and various other higher alcohols or corresponding

It is also known that if a substantially 25 equimolecular mixture of alcohol and hydrogen is passed over certain catalysts consisting of magnesium oxide mixed with other oxides and heated at between 220° and 350° C. under atmospheric pressure, there is a formation of higher alcohole more expecially of buttl alcohol. alcohols, more especially of butyl alcohol and hexyl alcohols, in addition to small quantities of acetaldehyde, butyraldehyde and crotonaldehyde.

The present invention has for its chief object a process which enables the conversion of alcohol almost exclusively into ethyl acetate to be effected. It is based upon the following observations:-

When alcohol vapours are passed, without special precautions, over a dehydro-genation catalyst which has been heated to a temperature between 200 and 260° C., to a temperature between 200 and 250 °C., there is observed simply a brisk evolution 45 of hydrogen with the production of acctaldehyde. However, if, instead of allowing the gases formed in the reaction to escape at the exit from the furnace, they are, after partial condensation, taken they to the furnace, with the side of some 50 back to the furnace, with the aid of some mechanical means, and forced to pass rapidly therethrough, it being arranged that the supply of gases rich in hydrogen

that are thus brought back over the catalyst is great in relation to the alcohol 55 supplied, the surprising fact is observed that a part of the alcohol is converted into ethyl acetate.

There is at the same time the production of a small quantity of acetaldehyde. 60 Now, it has also been found that the acetaldehyde itself can be converted, in turn, into ethyl acetate in the catalyst furnace under the conditions in which the principal reaction is effected.

Everything takes place as if the two

following reactions were produced: -

2CH, CH2OH__CH, COOCH2.CH3+2H2 2CH, CHO____ __CH_.COOCH_.CH_.

Under these conditions, it is therefore 70 recommended to separate from the reaction product the acetaldehyde and the alcohol which have not reacted and to cause them to re-enter the circuit with a view to their subsequent and total conversion into ethyl 75

The process forming the subject matter of the invention, which is based on these observations, therefore, consists essentially in passing the alcohol that is to be 80 converted over a dehydrogenation catalyst that is heated at between 200° and 260° C., subjecting the products rich in hydrogen that result from the catalytic conversion to a partial condensation and passing back 85 continuously over the catalyst, at the same vapours that have escaped condensation; preferably the inflow of alcohol and the return of the gases and vapours are regu- 90 lated in such a manner that the supply of the latter is great in relation to the supply of alcohol. In addition the acetaldehyde and the alcohol that have not reacted are recovered in the condensed 95

products and passed back over the catalyst.

It is to be noted that if a dilute alcohol is employed for effecting the reaction, the quantity of ester formed decreases rapidly whilst, at the same time, the production 100 of a large proportion of acetic acid is observed. It is therefore necessary to work with an anhydrous alcohol for the

purpose of obtaining the maximum rate of conversion into ester at the same time as a high yield and of avoiding the pro-duction of fatty acids having the same 5 number of carbon atoms as the alcohol treated.

There may be used as catalysts all the reduced metals that have a dehydrogenating action, provided that they pro-10 duce as little destruction as possible of the alcohol with formation of carbon monoxide, carbonic acid and hydro-carbons. It is nearly always advantageous to add a small quantity of promoter 15 oxides, among which the most interesting are the oxides of titanium, thorium and uranium.

The process is applicable not only to ordinary alcohol but also to other primary 20 aliphatic alcohols, the latter giving rise to corresponding esters possessing double

the number of carbon atoms.

The following description with reference to the accompanying drawing, which is given by way of example and represents, in its single Figure, the diagram of an apparatus that is suitable for carrying out the invention, will enable one to understand well how the latter may be carried 30 into effect:-

Anhydrous ethyl alcohol is passed, at the rate of 100 kilogrammes per hour, into a vaporiser 10 connected to a furnace 1 that contains a catalyst consisting reduced copper carried on infusorial earth and 10 per cent of uranium oxide as promoter. The catalyst being heated at 220° C., there is first of all observed a brisk evolution of hydrogen with the prodo duction of pure acetaldehyde which is partially condensed in a condenser 2 at the same time as the unconverted alcohol, the whole of the condensate being collected

in a vessel 3.

If, at this moment, the gases are taken off through a pipe 4 by means of a pump 5 and passed back over the catalyst along with the vapours of the alcohol the rate of supply of which remains unchanged, it 50 is observed that the liquids condensed in the vessel 3 contain, after passage through the catalyst furnace, a large quantity of ethyl acetate whilst, at the same time, the percentage of acetaldehyde is only slight. The acetaldehyde can be separated by distillation in a column 6 and brought

back continuously through a pipe 11 with the alcohol supplied. From that moment onwards the apparatus works normally.
It is arranged that the flow of gases that are rich in hydrogen and are brought back

over the catalyst should be high. For example, the action of the fan may be regulated to ensure a supply of 250 cubic 65 metres of gases per hour when the total

amount of alcohol supplied is 100 kilogrammes.

There escapes from the apparatus at 7, through a valve of any suitable type, hydrogen in a quantity corresponding to 70 the ethyl acetate that is formed. From the gases leaving the catalyst furnace, there is condensed in the vessel 3 an alcoholic liquid having approximately the following composition:-Ethyl acetate -- 17 per cent.

Acetaldehyde -1.5 0.15 ,, Acetic acid Butyl alcohol and higher alcohols -0.780

Water traces. Alcohol about 80.0

Various other substances to make up 100%.
The yield of ethyl acetate, in relation to the alcohol that has disappeared in the 85 reaction, is therefore of the order of 95 per cent. This ethyl acetate is separated per cent. This ethyl acetate is separated at the top of a distilling column 8, mixed with small quantities of alcohol and water, this mixture having a composition 90 that is intermediate between that of the ternary azeotropic mixture boiling at 70.3° C. and that of the binary esteralcohol azeotropic mixture boiling at 71.8° C. From this crude product, pure an- 95 hydrous ethyl acetate can be extracted by known methods.

The unconverted alcohol is withdrawn laterally from the distilling column 8 through the pipe 9 and is brought back 100 into the tank 13 with the supply of alcohol. The heavy products are withdrawn at 12 at the base of the column.

Finally, it is pointed out that the whole or a part of the process can be 105 carried out under a pressure that differs from the atmospheric pressure, if this measure is found to be convenient or advantageous.

Having now particularly described and 110 ascertained the nature of our said invention and in what manner the same is to be performed, we declare that what we

claim is:-1. Process for the conversion in a de- 115 hydrogenation reaction of an aliphatic primary alcohol into an ester containing double the number of carbon atoms com-prising essentially dehydrogenating the alcohol to be converted over a catalyst 120 consisting essentially of a reduced metal, said catalyst being maintained at a tem-perature between 200 and 260°, removing by condensation some of the gas-vapour product rich in hydrogen resulting from 125 the reaction and returning the gaseous remainder rich in hydrogen continuously to the catalysing zone to join with further fresh alcohol.

2. A process according to claim 1, in 130

which the catalyst consisting essentially of a reduced metal is activated by a promoter oxide or promoter oxides such, for example, as an oxide or oxides of 5 titanium, thorium or uranium.

3. Process according to claim 1 or claim 2, in which the alcohol to be converted is used in a substantially an-

hydrous state.

4. Process according to claim 1, 2 or 3, including the step of regulating the rate of admission of the returned gases so as

to be great relatively to the fresh alcohol.

5. Process according to claim 1 or
15 claim 2, including the step of separating
the resultant aldehyde and unconverted
alcohol in the condensate from the
resultant ester, and returning said aldehyde and unconverted alcohol to the 20 catalysing zone.

6. Process according to any of the preceding claims in which the alcohol employed is ethyl alcohol to obtain ethyl

acetate.

7. The process of conversion of aliphatic 25 primary alcohols into esters containing double the number of carbon atoms substantially as hereinbefore described.

8. The apparatus substantially as hereinbefore described with reference to the 30 accompanying drawing whenever employed for carrying out the process claimed in any of the preceding claims.

9. Organic esters whenever produced substantially as claimed in any of the 35 preceding claims 1 to 7.

Dated this 20th day of February, 1936.

For USINES DE MELLE.
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17. John Street, New York, U.S.A.
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